Novel Methylated Biomarkers and a Robust Assay to Detect Circulating Tumor DNA in Metastatic Breast Cancer

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Abstract

The ability to consistently detect cell-free tumor-specific DNA in peripheral blood of patients with metastatic breast cancer provides the opportunity to detect changes in tumor burden and to monitor response to treatment. We developed cMethDNA, a quantitative multiplexed methylation-specific PCR assay for a panel of ten genes, consisting of novel and known breast cancer hypermethylated markers identified by mining our previously reported study of DNA methylation patterns in breast tissue (103 cancer, 21 normal on the Illumina HumanMethylation27 Beadchip) and then validating the 10-gene panel in The Cancer Genome Atlas project breast cancer methylome database. For cMethDNA, a fixed physiologic level (50 copies) of artificially constructed, standard nonhuman reference DNA specific for each gene is introduced in a constant volume of serum (300 μL) before purification of the DNA, facilitating a sensitive, specific, robust, and quantitative assay of tumor DNA, with broad dynamic range. Cancer-specific methylated DNA was detected in training (28 normal, 24 cancer) and test (27 normal, 33 cancer) sets of recurrent stage IV patient sera with a sensitivity of 91% and a specificity of 96% in the test set. In a pilot study, cMethDNA assay faithfully reflected patient response to chemotherapy (N = 29). A core methylation signature present in the primary breast cancer was retained in serum and metastatic tissues collected at autopsy two to 11 years after diagnosis of the disease. Together, our data suggest that the cMethDNA assay can detect advanced breast cancer, and monitor tumor burden and treatment response in women with metastatic breast cancer. Cancer Res; 74(8); 2160–70. ©2014 AACR.

Introduction

Recent developments have shown that circulating cell-free DNA in blood offers a very convenient, noninvasive, and repeatable "liquid biopsy," thus providing a reliable template for assessing molecular markers that reflect tumor burden. Several DNA methylation studies have suggested clinical prognostic and/or predictive utility of methylated biomarkers for predicting outcome and monitoring treatment of breast cancer. DNA methylation is one of the earliest, most robust, and frequent alteration in cancer, leading to silencing of expression of tumor suppressor genes (1). Methylation of a tumor suppressor gene allele can act as an inactivating "hit," having a similar function as a genetic mutation or deletion. For examples, BRCA1 (breast, ovarian), PTEN (ovarian), HRK (prostate), APC (colorectal), and RASSF1A (multiple cancer cell types) have been found both methylated and mutated in cancer. Methylated RASSF1A and APC, identified in serum DNA from patients with breast cancer, were associated with a worse outcome (2). RASSF1A and NEUROD1 were shown to be useful for monitoring the efficacy of adjuvant therapy in patients with breast cancer (3, 4). Other biomarkers such as p16INK4A, CDH1, DAPK1, HIC1, RARB, CDH13, ESRI, and GSTPI, have been evaluated alone or in combination in serum (reviewed in ref. 5). However, a reproducible blood-based test for hypermethylated genes for diagnosis and follow-up of breast cancer has yet to be incorporated into a clinical laboratory assay. Discovery of new markers, as well as improvements in existing technologies, are needed to provide more robust, reproducible, quantitative, sensitive, and specific assays.

Here, we report the identification of novel methylated breast cancer genes as well as a robust serum/plasma-specific modified quantitative multiplexed methylation-specific PCR (QM- MSP) assay called cMethDNA, which enhances the methylation signal, while maintaining the high sensitivity, specificity, reproducibility, dynamic range, and quantitative advantages of the
standard QM-MSP assay (6–10). We balanced several criteria to select ten biomarkers that were simultaneously (i) highly and frequently methylated in breast tumor tissues (11), and in serum from patients with metastatic breast cancer and (ii) methylated at low levels in cell-free circulating serum DNA in normal individuals. Following validation of the 10-gene panel in The Cancer Genome Atlas project (TCGA) breast cancer methylome database, we developed and tested serum-specific prediction models, in test and training sets of sera from stage IV metastatic breast cancer. cMethDNA identified 91% of sera in patients with recurrent metastatic breast cancers with a specificity of 96% (area under the curve; AUC = 0.994; P < 0.0001) in the test set. Methylation levels reflected response to treatment, and circulating tumor DNA revealed a similar pattern of methylation as the solid tumor. We conclude that the cMethDNA assay performed on the 10-gene panel shows great potential for development as a clinical laboratory test for monitoring therapy and disease progression/recurrence.

Patients and Methods

Patients and sample collections

Whole blood and tissue were collected prospectively from women with stage IV metastatic breast carcinoma following disease recurrence after prior therapy [training set: J0214, NCT00080665; ref. 12; and J0425, NCT00274768 collected at JH 2004–2008; test set: J0524, Translational Breast Cancer Research Consortium (TBCRC) 005 collected 2004–2012], as well as from healthy controls (randomly divided into training and test sets: TBCRC 005). To evaluate concordance between sera and tissue, tissues were obtained from a subset of patients with cancer in the training set, along with additional samples collected at diagnosis of de novo metastatic disease (TBCRC 013 collected at participating institutions from 11/2009–4/2012). All the samples were collected with appropriate approval from the various Institutional Review Boards. Supplementary Materials and Methods has additional details. Patient characteristics are summarized in Table 1 and Supplementary Table S1.

Purification of cell-free circulating DNA

DNA extraction from serum was tested using three different serum DNA purification kits: QiaAmp MinElute Virus Spin Kit (ME; Qiagen), QiaAmp UltraSens Virus Kit (US; Qiagen), and Quick-gDNA MiniPrep (ZR; Zymo Research). The MinElute method was then selected for use in the study because of superior performance. External recombinant gene-specific standards (STDgene; 50 copies per gene for up to 12 genes) and carrier DNA/RNA (250 ng salmon sperm (Life Technologies), 250 ng tRNA (Roche Applied Science), and 5.6 μg “Carrier RNA” (Qiagen)) were added to each serum sample (300 μL), and cell-free DNA was extracted as per manufacturer’s instructions. Extracted DNA was then modified with sodium bisulfite, cleaned and eluted in 15 μL of water according to the EZ DNA Methylation Kit protocol (Zymo Research). Supplementary Materials and Methods have detailed protocols.

Extraction of tissue DNA

Tissue DNA was extracted overnight at 56°C in buffer containing TNES (10 mmol/L Tris, 150 mmol/L NaCl, 2 mmol/L EDTA, 0.5% SDS) and protease K (30 μg), heat inactivated 10 minutes at 90°C, then treated with sodium bisulfite as described previously (11, 13).

DNA methylation array

Details are presented in Supplementary Materials and Methods.

The cMethDNA assay

cMethDNA was performed in two sequential PCR reactions (Fig. 1A).

PCR reaction #1 (multiplex reaction). DNA was isolated from 300 μL serum containing 50 copies of each STDgene DNA, bisulfite treated, and amplified with a cocktail of external DNA-specific primer pairs specific for each of the 10 genes. Reaction #1 was then diluted 1:500–1:50,000.

PCR reaction #2 (real-time qMSP). Diluted amplicons (4 μL) from reaction #1 were amplified by real-time PCR, with the TARGETgene and reference STDgene in the same well. Patient samples were purified, assayed in duplicate, and results averaged. Detailed information is available in Supplementary Materials and Methods.

Calculation of methylation

For an individual serum sample, cMethDNA calculations were as follows: methylation index = [Methylated TARGETgene copies/(Methylated TARGETgene + STDgene) copies] (100); and cumulative methylation index (CMI) = the sum of all methylation index values within the gene panel. Serum samples were assayed in duplicate and then results were averaged.

For an individual sample, QM-MSP calculations: % methylation (%M) = [methylated TARGETgene copies/(methylated TARGETgene + unmethylated TARGETgene) copies](100); CMI = the sum of all %M values within the panel.

Statistical analysis

cMethDNA data analyses were performed using GraphPad Prism version 5.0 (GraphPad Software), SAS software (v 9.2, SAS Institute Inc.), or with R version 2.15.2 (October 26, 2012). Statistical tests were two sided and considered statistically significant at P < 0.05 unless otherwise stated. Distributions of cMethDNA data between independent groups were described using box plots and difference was tested using nonparametric Mann–Whitney test. The Wilcoxon signed rank test was performed when comparing two related samples (e.g., measurements on the same subjects). Receiver operating characteristic (ROC) analyses were used to characterize performance and define laboratory thresholds. The performance of the 10-gene panel was characterized through estimating the area under the ROC curve, sensitivity, specificity, classification accuracy, and likelihood ratio along with the 95% confidence intervals (CI). Details of statistical analyses can be found in Supplementary Materials and Methods.

Results

Whole-genome methylation array

For the selection of the 10-gene panel used in the cMethDNA assay, we relied heavily on our previously published whole-genome analysis of breast tissue DNA (N = 103 tumors, N = 21...
normal breast samples, Infinium Human Methylation27K Beadchip; ref. 11), along with data generated in a new array analysis of DNA methylation in serum \[N = 6\] recurrent metastatic breast cancer sera, \[N = 5\] normal sera, \[N = 4\] leukocyte pools (5 normal individuals per pool). For identifying candidate markers in primary breast tissue, we serially selected: (i) 8,376 of 27,578 probes with \(SD > 0.100\) between tumor tissues and probe detection \(P < 0.0001\), (ii) 2,674 of 8,376 probes having at least 1.5-fold higher mean methylation in tumors than in normal breast organoids, and (iii) 1,752 of 2,674 probes with \(\beta\)-methylation \(< 0.15\) in normal breast tissue adjacent to tumor (\(N = 6\), laser microdissected). We further filtered this pool of candidate markers on the basis of data generated in sera, choosing, (iv) 212 of 1,752 probes \(\geq 2.0\)-fold more methylated in cancer serum than in normal serum, and 198 of 212 probes with \(\beta\)-methylation \(< 0.15\) in individual serum samples.

The final 10-gene marker panel was chosen by careful inspection of the 198 probes, prioritizing probes that combined very low methylation in normal samples with frequent and high levels of methylation in both estrogen receptor-positive (ER\(^+\)) and ER-negative cancers. Eighteen of these probes were

### Table 1. Patient characteristics

<table>
<thead>
<tr>
<th>Patient characteristics - metastatic breast cancer</th>
<th>Recurrent Training set</th>
<th>Recurrent Test set</th>
<th>De novo TBCRC 013</th>
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<tr>
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<td><strong>n = 33</strong></td>
<td><strong>n = 18</strong></td>
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<tr>
<td>Race</td>
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<td></td>
</tr>
<tr>
<td>Caucasian</td>
<td>17</td>
<td>23</td>
<td>13(^a)</td>
</tr>
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<td>6</td>
<td>10</td>
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<td>3</td>
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<tr>
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<td>20</td>
<td>7</td>
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<tr>
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<td></td>
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</tr>
<tr>
<td>ER/PR(^+), HER2 negative</td>
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<td>17</td>
<td>9</td>
</tr>
<tr>
<td>HER2 positive</td>
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<td>10</td>
<td>8</td>
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<td>Triple negative (ER, PR, HER2 negative)</td>
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<td><strong>n = 28</strong></td>
<td><strong>n = 27</strong></td>
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<tr>
<td>Patient characteristics</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Race</td>
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<td></td>
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<tr>
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<td>12</td>
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</tr>
<tr>
<td>Black</td>
<td>11</td>
<td>15</td>
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<tr>
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<tr>
<td>Range</td>
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<td>20–63</td>
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\(^a\)Includes one Hispanic, 12 non-Hispanic.
present among a set of 100-candidate recurrence markers identified previously in the tissue DNA methylation array (11). Among these were seven novel markers (AKR1B1, COL6A2, GPX7, HIST1H3C, HOXB4, RASGRF2, and TM6SF1) and two known ones, ARHGEF7 and TMEFF2. RASSF1 was selected to complete the 10-gene marker panel (Supplementary Table S2; Supplementary Figs. S1A and S2).

The 10-gene methylation panel was verified in silico for sensitivity and specificity in TCGA Project databases (N = 316 breast cancer, N = 27 normal breast samples, BRCA; Supplementary Fig. S1B), and subsequently verified in sera using the cMethDNA assay. The 10-probe test panel outperformed 97.9% of 100,000 iterations of randomly created 10-probe panels drawn from the TCGA database (Supplementary Fig. S1C).

cMethDNA assay

The cMethDNA assay is a refinement of the QM-MSP method used extensively by our laboratory and others (6, 7, 9, 10, 14). Here, a low fixed physiologic level of recombinant gene-specific standard reference DNA is introduced into a constant volume of serum (50 copies of gene-specific standard DNA at 300 µL serum) before purification of the DNA. This fixes a relatively high constant ratio of methylated DNA to reference DNA, which is quantified after multiplex and quantitative real-time PCR. Primer/probe sequences are in Supplementary Table S3.

Technical validation of the cMethDNA assay

Intra-assay testing. To directly compare the QM-MSP and cMethDNA assays, DNA from recurrent metastatic breast cancer patient sera was tested by both methods for the

![Figure 1. The cMethDNA assay.](image)
RASSF1A gene (Fig. 1B). The robustness of methylation values and frequency of detection of hypermethylated RASSF1A were higher with cMethDNA compared with the QM-MSP assay \( P = 0.008 \) (Wilcoxon signed rank test). In a second experiment, six aliquots of serum from a single normal donor (300 μL serum per assay point) were spiked with a physiologic range (0, 50, 200, 800, or 3,200 copies) of fully methylated DNA from the breast cancer cell line, MDA-MB-453. DNA was purified, multiplexed, and tested by both cMethDNA and QM-MSP methods for the 10-gene panel (excluding HIST1H3C, which is not methylated in this cell line; Fig. 1C). Again, the cMethDNA assay reported significantly higher methylation than the QM-MSP method at all levels (50 copies, \( P = 0.008 \); 200 copies, \( P = 0.006 \); 800 copies, \( P = 0.006 \); 3,200 copies, \( P = 0.004 \); one-sided \( t \) test, permuted; Fig. 1C).

**Reproducibility of replicates, comparison of extraction methods.** We performed a series of tests to establish linearity and sensitivity of the cMethDNA assay. Three different serum DNA extraction methods were evaluated (MinElute Virus Kit, Qiagen UltraSens Virus Kit, Qiagen, and Quick-gDNA Prep, Zymo Research). Methylated DNA was spiked as described above (Fig. 1C); six replicates per point per extraction method were tested by cMethDNA (Fig. 2A–C). The MinElute Virus Spin Kit method showed the highest reproducibility and the smallest interassay coefficients of variation (CV = 29%, 12%, 4.6%, 2.5%, for 50, 200, 800, or 3,200 methylated DNA copies, respectively; Supplementary Table S4).

**Interuser reproducibility.** The ability for two individuals to perform the cMethDNA assay and to arrive at similar results is important. For the 10-gene panel cumulative methylation levels obtained by two users were compared after independently extracting DNA and performing the cMethDNA assay on duplicate aliquots of patient sera \( (N = 13) \). The interuser reproducibility, tested for intraclass correlation coefficient (ICC) showed strong agreement between users \( (ICC = 0.99; 95\% \text{ CI}, 0.96–1.00; \text{Fig. 2D}) \).

In summary, the cMethDNA approach results in an enhanced methylation signal, minimizes the effects of technical variability in purification of DNA, and because it uses spiked DNA as a standard, is not affected by potential day-to-day fluctuations in total serum DNA content that may be independent of changes in the tumor burden.

**Detection of methylated DNA in serum of patients with metastatic breast cancer**

**Assay specificity.** To verify the performance of the 10-gene marker panel in the cMethDNA assay for detection of circulating cell-free tumor DNA, we evaluated independent training and test sets of sera collected in prospective clinical trials of patients with recurrent stage IV breast cancer conducted at Johns Hopkins (Baltimore, MD) and through the TBCRC. The training set consisted of 52 serum samples (28 normal and 24 cancer). Sera from patients with cancer had significantly higher levels of methylated genes than women without cancer.

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**Figure 2.** cMethDNA assay validation. A–C, three serum purification methods (Qiagen MinElute, Qiagen Ultra Sens, and Zymo Research Quick g-DNA) were tested in conjunction with the cMethDNA method in normal serum aliquots spiked with 0 to 3,200 methylated copies, prepared from one master stock. Replicates were purified and multiplexed separately. Box-whisker plots show the median and full range of CMI (y-axis) for replicates of each sample (x-axis). Statistical significance (Mann–Whitney test) is indicated by \( P \) values. The % CV (a normalized measure of frequency distribution) is shown for each test in Supplementary Table S4. D, interuser reproducibility was evaluated for a set of thirteen patient serum samples processed independently by two investigators. User performance was evaluated for the 10-gene panel \( (ICC = 0.99; 95\% \text{ CI}, 0.96–1.00) \).
(median CMI = 117.3 and 0.04, respectively; \( P < 0.0001 \), Mann–Whitney test; Fig. 3A and B; Table 2A). ROC analyses identified a threshold of 6.9 units that maximized the sum of sensitivity and specificity in the training set (Table 2B). The assay specificity was 96.4% (27/28; 95% CI, 81.7–99.9%) when sensitivity was 91.7% (22/24; 95% CI, 73.0–99.0%) for a likelihood ratio = 25.7 and an overall classification rate of 94% (49/52; 95% CI, 84–99%). The ROC analysis AUC = 0.95 (95% CI, 0.87–1.02; \( P < 0.0001 \)). These findings were further verified in a test set of patient samples (27 normal, 33 cancer; Table 1 and Supplementary Table S1A). Consistent with findings in the training set, cumulative methylation levels were significantly higher in patients with metastatic breast cancer, compared with women without cancer (\( P < 0.0001 \), Mann–Whitney test; Fig. 3A and B; Table 2A). Using the classification rule derived on the training data, the observed assay specificity was 100% (27/27; 95% CI, 87.2–100%) at a sensitivity of 90.9% (30/33; 95% CI, 75.7–98.1%) for an overall classification accuracy of 95% (57/60; 95% CI, 86–99%) and a likelihood ratio = 24.6 (Table 2B). As in the training set, the median frequency of methylation of most genes was high (median = 38%, range = 18%–67%; Fig. 3C; Table 2C). RASSF1A had the highest incidence of methylation among all genes in both training and test sets (75% and 67%, respectively). Individually RASSF1A, HIST1H3C, RASGRF2,
Table 2. cMethDNA ability to detect circulating cell-free tumor DNA

<table>
<thead>
<tr>
<th>CMI (10 genes) in cancer and normal sera</th>
<th>Training set</th>
<th>Test set</th>
</tr>
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<tbody>
<tr>
<td>N</td>
<td>Normal</td>
<td>Stage IV</td>
</tr>
<tr>
<td>Minimum</td>
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<tr>
<td>25% percentile</td>
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<tr>
<td>Median</td>
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<tr>
<td>75% percentile</td>
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</tr>
<tr>
<td>Maximum</td>
<td>7</td>
<td>616</td>
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<tr>
<td>Mean</td>
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<td>165</td>
</tr>
<tr>
<td>Lower 95% CI of mean</td>
<td>0</td>
<td>99</td>
</tr>
<tr>
<td>Upper 95% CI of mean</td>
<td>2</td>
<td>230</td>
</tr>
</tbody>
</table>

ROC analysis

- Area under the curve: Training set = 0.950, Test set = 0.994
- 95% CI: Training set = 0.874–1.027, Test set = 0.984–1.005
- P < 0.0001
- Sensitivity: Training set = 92%, Test set = 91%
- Specificity: Training set = 96%, Test set = 100%
- Likelihood ratio: Training set = 25.7, Test set = 24.6
- Classification accuracy: Training set = 94%, Test set = 95%

Methylation frequency

<table>
<thead>
<tr>
<th>Gene</th>
<th>Training set Cancer (n = 24)</th>
<th>Control (n = 28)</th>
<th>Test set Cancer (n = 33)</th>
<th>Control (n = 27)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AKR1B1</td>
<td>15 (63%)</td>
<td>0</td>
<td>14 (42%)</td>
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</tr>
<tr>
<td>ARHGEF7</td>
<td>7 (29%)</td>
<td>0</td>
<td>6 (18%)</td>
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<tr>
<td>COL6A2</td>
<td>9 (37%)</td>
<td>1 (4%)</td>
<td>12 (36%)</td>
<td>0</td>
</tr>
<tr>
<td>GPX7</td>
<td>5 (21%)</td>
<td>0</td>
<td>10 (30%)</td>
<td>0</td>
</tr>
<tr>
<td>HOXB4</td>
<td>11 (46%)</td>
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<td>12 (36%)</td>
<td>0</td>
</tr>
<tr>
<td>RASGRF2</td>
<td>12 (50%)</td>
<td>2 (7%)</td>
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<td>RASSF1A</td>
<td>18 (75%)</td>
<td>1 (4%)</td>
<td>22 (67%)</td>
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<tr>
<td>TM6SF1</td>
<td>10 (42%)</td>
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<td>12 (50%)</td>
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</table>

\(^a\) Positive methylation defined as CMI > 7.0 (set by ROC).
\(^b\) Positive methylation defined as > 1 unit of methylation index.
COL6A2, HOXB4, and AKR1B1 genes had the strongest performance in the cMethDNA assay (P = 0.0003 to P < 0.0001, Mann–Whitney test of training set; Fig. 3C; Table 2C). A 6-gene panel of the six most robust biomarkers performed nearly as well as the 10-gene panel (Supplementary Fig. S3). In normal samples, no significant age-dependent changes in cumulative methylation were observed (N = 55; one-way ANOVA, P = 0.8988 for quartiles Table 1 and Supplementary Fig. S4).

Utility of cMethDNA to monitor response to treatment

To determine whether the cMethDNA assay might be used effectively to monitor response to chemotherapy, we evaluated sera collected from patients with metastatic breast cancer who participated in two prospective clinical trials J0214 and J0425 (15). Serum samples (N = 58) were collected from 29 patients at baseline before initiation of treatment, and 18 to 49 days (median 21 days) after initiation of a new chemotherapy regimen (Supplementary Table S1B). The results showed a statistically significant decrease in median serum DNA methylation levels in patients having stable disease (SD) or with a therapeutic response (e.g., partial response; PR) according to Response Evaluation Criteria in Solid Tumors (RECIST) criteria (P = 0.010; Fig. 4A). This decrease in methylation was not observed in patients with progressive disease (PD; P = 0.659; Wilcoxon signed rank test; Fig. 4A).

Because a single time point posttherapy may have limited potential to predict outcome as assessed by imaging several weeks to months later, we evaluated a subset of patients (13 of 29 with sera collected at three or more time points (total of 54 sera)) during different cycles of the same therapy (Fig. 4B–G; Supplementary Fig. S5 and Supplementary Table S1B). Ten of thirteen patient sera showed methylation levels reflective of decreases in tumor burden during stable disease or PR, and increasing levels of methylation during progressive disease as defined by RECIST criteria. These data show the potential of cMethDNA to reveal therapeutic response at an early time point during treatment and provide important information for clinicians that could aid decision making about further therapy.

Patterns of methylation are retained between primary tumor and metastasis

Previously, we observed similarity in the biomarker expression and methylation profiles of primary breast cancer and distant metastases collected from individuals within 4 hours of death (N = 10; 2–11 years after first diagnosis of breast cancer; age 33–79 years ref. 16). With the 10-gene marker panel, a striking concordance was observed between the methylated gene patterns in the samples from primary, metastases, and serum of the same patient (Fig. 5). In this analysis, serum was quantified with the cMethDNA assay, and tissues were quantified with the QM-MSP assay. Results are plotted on the same y-axis, although cMethDNA is more robust. Comparative analyses indicated >50% of individual patient serum/tissue pairs or tissue/tissue pairs were concordant for ≥9 markers, and all had ≥6 matches (Supplementary Fig. S6). Because by chance alone, the median number of matches predicted is 7 (range = 3–9), based on the overall frequency of the genes evaluated, results indicate that methylation profiles were more patient specific than random. Concordance between serum and primary and/or metastases was also observed in two other independent series of samples collected in clinical trials: (i) a subset of patients in the study training set (10 of 24 patients; Table 1 and Supplementary Table SIC and Supplementary Fig. S7A) and (ii) from untreated patients diagnosed with de novo metastatic breast cancer (18 patients; Table 1 and Supplementary Table SIC, in TBCRC 013, and Supplementary Fig. S7B). As in the autopsy study, consistent profiles of hypermethylation were observed between the primary tumor, serum, and distant metastasis samples, even though sera were tested with the cMethDNA method and tissues by the QM-MSP method.

Collectively, these studies provide proof-of-principle that a core pattern of methylation is retained in the distant metastasis and serum of a given patient over time, perhaps decades.

Utility of the 10-gene panel to detect other tumor types

Careful analysis and selection of breast cancer markers determined in tissues allowed us to build the 10-gene panel, which was later confirmed in serum DNA from patients with metastatic breast cancer and validated in the independent TCGA breast cancer methylation tissue DNA array database. To determine whether an in silico analysis of other cancer tissue databases will indicate the potential use of the 10-gene panel for detecting circulating methylated markers in other types of cancer, we examined several Infinium HumanMethylation27 array databases from TCGA (Supplementary Fig. S8A–S8F). On the basis of ROC analyses, compared with breast (BRCA, AUC = 0.950), the panel performed with a high level of efficiency in three other tumor types, including lung (LUNG, AUC = 0.969), colon (COAD, AUC = 0.995), and rectum (READ, AUC = 0.997). However, the panel displayed poor performance with other tumor types, such as ovarian (OV, AUC = 0.668), kidney (KIRC, AUC = 0.725), and stomach (STAD, AUC = 0.792). Thus, the panel, similar to breast carcinomas, shows specificity for the highly prevalent lung and colorectal adenocarcinomas and may not be useful for others such as ovarian, kidney, and stomach cancers.

Discussion

We have derived an informative panel of novel gene markers for detecting tumor-specific circulating, cell-free, methylated DNA in the sera of patients with metastatic breast cancer. Using this panel, we have demonstrated broad dynamic range, reproducibility, sensitivity, specificity, and accuracy of the cMethDNA assay for detecting methylated DNA in the vast majority of patients with newly diagnosed and recurrent stage IV breast cancer. We have also provided preliminary evidence for the potential of the cMethDNA assay to aid in monitoring disease during chemotherapy, and shown that a core methylation pattern typical of each primary tumor is retained in the metastatic lesions and serum over a long period of time (2–11 years). TCGA tumor tissue methylation arrays revealed that this panel of ten markers is methylated and likely to perform well to detect a variety of tumor types (i.e., lung, colon, rectal, and possibly uterine carcinomas and glioblastomas) and would have less utility for other tumors (i.e., ovarian, kidney,
stomach). Shortly, planned studies will test the prognostic and predictive utility of this assay in an external sample set from larger prospective clinical trials.

The challenge to developing a reliable blood test for cancer has been finding the virtual “needle in the haystack” or desired methylated marker in the vast excess of unmethylated normal DNA. Our new cMethDNA assay overcomes this barrier by incorporating two innovations: first, spiking of serum with physiologic levels of recombinant standards matched to the gene of interest by size and homology at 5′ and 3′ ends that provide an internal reference for recovery and quantification of target DNA. Second, the inclusion of a multiplexed, preamplification step increases the dynamic range of detectability of the target gene and enables quantitation of a panel of markers in the amount of DNA normally used to evaluate one gene. The use of external standards has a further advantage in that it avoids the pitfalls inherent to endogenous reference DNAs, where total DNA may fluctuate up to 15-fold under healthy conditions.

**Figure 4.** Monitoring response to treatment. A, cMethDNA was performed on serum samples obtained from patients with metastatic breast cancer at baseline (pre) and after 18 to 49 days of therapy (post). Data are plotted according to CMI in individual patients judged by RECIST criteria to have stable/responsive disease or progressive disease after 8 to 12 weeks. Pre- and post-difference in CMI was evaluated using Wilcoxon signed rank test. B–G, representative plots of CMI of patient (ST#) sera assessed by cMethDNA during the course of treatment. Patients received either 28-day cycles of docetaxel or 21-day cycles of capecitabine, indicated by shading. C, cycles of treatment; Imaging- and RECIST criteria-assessed progressive disease.
independent of tumor burden, such as occurs in exercise over-training (17, 18), trauma (19), surgery (20), sepsis (21), chronic inflammatory diseases (22–24), and pregnancy (25, 26).

Features critical to the high sensitivity and specificity of the cMethDNA assay are the identification of biomarkers through whole-genome analysis of solid tumor and metastatic cancer sera, a careful filtration process, and the final selection of a panel of hypermethylated genes that were specifically and frequently methylated in breast cancer. Among the cancer-specific markers in our panel used for detecting cell-free DNA in serum by the cMethDNA assay and tissue DNA by QM-MSP, the HOXB4, RASGRF2, AKR1B1, TM6SF1, COL6A2, GPX7, HIST1H3C genes are being reported here for the first time in breast cancer. Interestingly, HOXB4 also hosts the regulatory region for miRNA (miR)-10a near its genome and is hypermethylated in 90% of hepatocellular carcinomas (27). In breast cancer, the cMethDNA assay had a classification accuracy of up to 95% that distinguished cancer versus normal sera with a sensitivity of more than 90% and specificity of nearly 100%, which suggest a potential for greater clinical utility compared with CA27.29 (28), carcinoembryonic antigen (CEA; ref. 29), circulating tumor cells (CTC; ref. 30), and other circulating tumor DNA tests (31, 32).

The application of cMethDNA as a noninvasive indicator of tumor burden and therapeutic response was tested using samples collected as secondary endpoints from three prospective clinical trials of women with metastatic breast cancer that was newly diagnosed or progressing after initial palliative chemotherapy. In a pilot study, the cMethDNA assay was able to detect serum methylation levels that reflected tumor burden based on RECIST criteria. In the majority of cases, changes in methylation were tracked with disease progression. Investigations of sera during subclinical stages of breast cancer recurrence (adjuvant surveillance) are also needed. Although imaging studies (in addition to clinical assessment) are the current standard to evaluate response after 8 to 12 weeks of initiating a new systemic, tests to monitor disease prognosis and predict therapy benefit earlier remain an unmet need. An ongoing trial, SWOG 0500, (NCT00382018) is evaluating the role of enumerating CTCs in this setting, but may be hampered by the low sensitivity of that commercial assay. The cMethDNA assay seems to be promising in this setting, and we are embarking on a comparative analysis of the prognostic and predictive utility of cMethDNA, CA27.29, CEA, and CTC, using prospectively collected serum to predict response to therapy in metastatic breast cancer. Investigations of sera obtained during the subclinical stages of breast cancer recurrence (adjuvant surveillance) are also needed.

In summary, the cMethDNA assay is a promising new liquid biopsy tool for detecting tumor-specific cell-free circulating DNA in a noninvasive manner. With further refinement, it could serve to monitor response to therapy, potentially prognosticate disease outcome, and serve as an early indicator of tumor recurrence.

Disclosure of Potential Conflicts of Interest
S. Sukumar and M.J. Fackler have ownership interest (including patents) in a patent application for the cMethDNA method (no financial interest) and Patent for RASSF1A with the QM-MSP method (no financial interest). Z. Lopez Bujanda has ownership interest (including patents) in a patent application for the cMethDNA method (no financial interest). W.W. Teo has ownership interest (including patents) in a cMethDNA patent application (no financial interest). S. Sukumar is a consultant/advisory board member of California Breast Cancer Research Foundation. No potential conflicts of interest were disclosed by the other authors.

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Conception and design of clinical trial only: L.A. Carey, A.C. Wolff
Development of methodology: M.J. Fackler, Z. Lopez Bujanda, W.W. Teo, A.C. Wolff, S. Sukumar
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References


Correction: Novel Methylated Biomarkers and a Robust Assay to Detect Circulating Tumor DNA in Metastatic Breast Cancer

In this article (Cancer Res 2014;74:2160–70), which appeared in the April 15, 2014, issue of Cancer Research (1), a symbol in the précis was incorrectly placed. The corrected précis is below. The publisher regrets this error.

The online version has been corrected and no longer matches the print.

This report describes a multiplexed PCR method for detecting circulating methylated DNA in the serum of patients with metastatic breast cancer, offering >90% sensitivity and specificity to detect disease.

Reference


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Novel Methylated Biomarkers and a Robust Assay to Detect Circulating Tumor DNA in Metastatic Breast Cancer

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